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(54) NOVEL VEGF-LIKE FACTORS

homology with a VEGF-C gene, a member of the VEGF family, has been isolated by the PCR method using primers designed based on the sequence of EST that is assumed to be homologous with the C-termial region of the VEGF-C gene. Mouse and rat genes have been isolated based on the human gene isolated as above. A protein encoded by the above human gene has been isolated by introducing the gene into *Escherichia coli* and expressing it. The isolated protein and genes can be applied to, for example, gene therapy for the VEGF-D deficiency, wound healing, and promotion of collateral vessel formation. Furthermore, VEGF-D protein inhibitors can be used as a novel anticancer drug, etc.

Description

Technical Field

5 [0001] The present invention relates to a protein factor involved in angiogenesis in humans and falls in the field of genetic engineering.

Background Art

10 [0002] The process of angiogenesis, in which endothelial cells existing in the inner wall of blood vessels of animals generate new blood vessels, is triggered by transduction of a specific signal. A variety of substances are reportedly involved in this signal transduction. The most notable substance among them is the vascular endothelial growth factor (VEGF). VEGF is a protein factor which was isolated and purified, and can increase the proliferation of endothelial cells and the permeability of blood vessels (Senger, D. R. et al., Science 219: 983-985 (1983); Ferrara, N. and Henzel, W. J., Biochem. Biophys. Res. Commun. 161: 851-858 (1989)). It has been reported that the human VEGF gene contains eight exons and produces four subtypes consisting of 121, 165, 189, or 206 amino acid residues, depending on the difference in splicing, which causes different secretionpatterns (Houck, K. A. et al., Mol. Endocrinol. 5: 1806-1814 (1991)). It has also been reported that there is a VEGF-specific receptor, fit-1, and that the binding of VEGF to flt-1 is important for the signal transduction (Vries, C. D. et al., Science 255: 989-991 (1992)).

[0003] Placental growth factor (PIGF) and platelet-derived growth factor (PDGF) have thus far been isolated and are factors related to VEGF. These factors are found to promote proliferation activities of vascular endothelial cells (Maglione, D. et al., Proc. Natl. Acad. Sci. USA 88: 9267-9271 (1991); Betsholtz, C. et al., Nature 320: 695-699 (1986)). In addition, VEGF-B (Olofsson, B. et al., Proc. Natl. Acad. Sci. USA 93: 2576-2581 (1996)) and VEGF-C (Lee, J. et al., Proc. Natl. Acad. Sci. USA 93: 1988-1992 (1996); Joukov, V. et al., EMBO J. 15, 290-299 (1996)) have recently been isolated.

[0004] These factors appear to constitute a family, and this may contain additional unknown factors.

[0005] It has been suggested that VEGF is involved in not only vascular formation at the developmental stage but also in the pathological neovascularization associated with diabetes, rheumatoid arthritis, retinopathy, and the growth of solid tumors. Furthermore, in addition to its vascular endothelial cell growth-promoting effects listed above, VEGF's ability to increase vascular permeability was suggested to be involved in the edema formation resulting from various causes. Also, these VEGF family factors may act on not only the blood vessels but also the blood cells and the lymphatic vessels. They may thus play a role in the differentiation and proliferation of blood cells and the formation of lymphatic vessels. Consequently, the VEGF family factors are presently drawing extraordinary attention for developing useful, novel drugs.

Disclosure of the Invention

[0006] An objective of the present invention is to isolate a novel protein belonging to the VEGF family and a gene encoding the protein. We searched for genes having homology to VEGF-C, which is a recently cloned VEGF family gene, against Expressed Sequence Tags (EST) and Sequence Tagged Sites (STS) in the GenBank database. As a result, we found an EST that was assumed to have homology to the C-terminal portion of VEGF-C. We then designed primers based on the sequence, and amplified and isolated the corresponding cDNA using the 5' RACE method and the 3' RACE method. The nucleotide sequence of the isolated cDNA was determined, and the deduced amino acid sequence therefrom revealed that the amino acid sequence had significant homology to that of VEGF-C. Based on the homology, we have assumed that the isolated human clone is a fourth member of the VEGF family (hereinafter designated as VEGF-D). We have also succeeded in expressing the protein encoded by the isolated human VEGF-D gene in E. coli cells, and have also purified and isolated it. Furthermore, we have succeeded in isolating the mouse and rat VEGF-D genes using the isolated human VEGF-D gene.

[0007] In particular, the present invention relates to a novel protein belonging to the VEGF family and a gene encoding

- the protein. More specifically it relates to
 - (1) A protein shown by SEQ ID NO.1 or having the amino acid sequence derived therefrom in which one or more amino acids are substituted, deleted, or added;
 - (2) A protein encoded by a DNA that hybridizes with the DNA shown by SEQ ID NO. 2;
- (3) A DNA encoding the protein of (1);

- (4) A DNA hybridizing with the DNA shown by SEQ ID NO. 2;
- (5) A vector containing the DNA of (3) or (4);
- (6) A transformant carrying the vector of (5);

- (7) A method of producing the protein of (1) or (2), which comprises culturing the transformant of (6);
- (8) An antibody binding to the protein of (1) or (2):

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- (9) A method of screening a compound binding to the protein of (1) or (2), which comprises a step of detecting the activity of the protein of (1) or (2) to bind to a test sample; and
- (10) A compound binding to the protein of (1) or (2), wherein said compound has been isolated by the method of (9).

[0008] The protein of the present invention (VEGF-D) has significant homology to VEGF-C and can be considered to be a fourth factor of the VEGF family. Since the major function of VEGF is vascular formation at the developmental stage and VEGF is considered to be involved in the pathological neovascularization associated with diabetes, rheumatoid arthritis, retinopathy, and the growth of solid tumors, the protein of the present invention is thought to have similar functions.

[0009] A person skilled in the art could prepare functionally equivalent proteins through modifying VEGF-D of the present invention by adding, deleting, or substituting one or more of the amino acids of VEGF-D shown by SEQ ID NO.

1 using known methods. Modifications of the protein can also occur naturally in addition to the artificial modifications described above. These modified proteins are also included in the present invention. Known methods for adding, deleting, or substituting amino acids include the overlap extension polymerase chain reaction (OE-PCR) method (Gene, 1989, 77 (1): 51).

[0010] The DNA encoding VEGF-D of the present invention, shown by SEQ ID NO. 2, is useful for isolating DNAs encoding the proteins having similar functions to VEGF-D in other organisms. For example, a person skilled in the art could routinely isolate homologs of human VEGF-D of the present invention from other organisms by allowing the DNA shown by SEQ ID NO. 2, or part thereof, as a probe, to hybridize with the DNA derived from other organisms. The DNA that hybridizes with the DNA shown by SEQ ID NO. 2 is also included in the present invention. The other organisms include mice, rats, and rabbits.

[0011] The DNA encoding a protein that is functionally equivalent to VEGF-D usually has high homology to the DNA shown by SEQ ID NO. 2. The high homology used herein means at least 70% or higher, more preferably 80% or higher, and still more preferably 90% or higher of sequence homology.

[0012] An example of the hybridization conditions for isolating the DNA having high homology will be given below. Prehybridization is performed in ExpressHyb Solution at 68°C for 30 minutes. The probe labeled with a radioisotope is denatured at 95°C to 100°C for 2 to 5 minutes and rapidly chilled on ice. The probe is added to a new ExpressHyb Solution. The blot is transferred to the solution containing the probe and allowed to hybridize under a temperature gradient of 68°C to 55°C for 2 hours. The blot is washed four times, for 10 minute each, with a 2 x SSC solution containing 0.05% SDS at room temperature. The blot is then washed with a 0.1 x SSC solution containing 0.1% SDS at 45°C for 3 minutes. The blot is subjected to autoradiography.

[0013] An example of the hybridization conditions for isolating the DNA having very high homology will be given below. Prehybridization is performed in ExpressHyb Solution at 68°C or 30 minutes. The probe labeled with a radioisotope is denatured at 95°C to 100°C for 2 to 5 minutes and rapidly chilled on ice. The probe is added into a new ExpressHyb Solution. The blot is transferred into the solution containing the probe, and allowed to hybridize at 68°C for 1 hour. The blot was washed four times, for 10 minute each, with a 2 x SSC solution containing 0.05% SDS at room temperature.

The blot was then washed with a 0.1 x SSC solution containing 0.100 Containing 0.05% SDS at room temperature.

The blot was then washed with a 0.1 X SSC solution containing 0.1% SDS at 50°C for 40 minutes, during which the solution was replaced once. The blot was then subjected to autoradiography.

[0014] Note that the hybridization condition can vary depending on the length of the probe (whether it is an oligomer or a probe with more than several hundred bases), the labeling method (whether the probe is radioisotopically labeled or non-radioisotopically labeled), and the type of the target gene to be cloned. A person skilled in the art would properly select the suitable hybridization conditions. In the present invention, it is especially desirable that the condition does not allow the probe to hybridize with the DNA encoding VEGF-C.

[0015] The DNA of the present invention is also used to produce VEGF-D of the present invention as a recombinant protein. Specifically, the recombinant protein can be produced in large quantity by incorporating the DNA encoding VEGF-D (for example, the DNA shown by SEQ ID NO. 2) into a suitable expression vector, introducing the resulting vector into a host, and culturing the transformant to allow the recombinant protein to be expressed.

[0016] The vector to be used for producing the recombinant protein is not particularly restricted. However, vectors such as pGEMEX-1 (Promega) or pEF-BOS (Nucleic Acids Res. 1990, 18(17): p.5322) are preferable.. Suitable examples of the host into which the vector is introduced include E. coli cells, CHO cells, and COS cells.

[0017] The VEGF-D protein expressed by the transformant can be purified by suitably combining purification treatments such as solubilization with a homogenizer or a sonicator, extraction by various buffers, solubilization or precipitation by acid or alkali, extraction or precipitation with organic solvents, salting out by ammonium sulfate and other agents, dialysis, ultrafiltration using membrane filters, gel filtration, ion exchange chromatography, reversed-phase chromatography, counter-current distribution chromatography, high-performance liquid chromatography, isoelectric

focusing, gel electrophoresis, or affinity chromatography in which antibodies or receptors are immobilized.

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[0018] Once the recombinant protein is obtained, antibodies against it can be prepared using known methods. The known methods include preparing polyclonal antibodies by immunizing rabbits, sheep, or other animals with the purified protein, and preparing monoclonal antibodies from the antibody-producing cells of immunized mice or rats. These antibodies will make it possible to quantify VEGF. Although the antibodies thus obtained can be used as they are, it will be more effective to use the humanized antibodies to reduce the immunogenicity. The methods of humanizing the antibodies include the CDR graft method and the method of directly producing a human antibody. In the CDR Graft method, the antibody gene is cloned from the monoclonal antibody-producing cells and its antigenic determinant portion is transplanted into an existing human antibody. In the method of directly producing a human antibody, a mouse whose immune system has been replaced by the human immune system is immunized, similar to ordinary monoclonal antibodies. The VEGF-D protein or its antibody thus obtained can be administered into the body by subcutaneous injection or a similar method.

[0019] A person skilled in the art could screen compounds that bind to the protein of the present invention by known methods.

[0020] For example, such compounds can be obtained by making a cDNA library on a phage vector (such as \(\lambda\)gt11 and ZAP) from the cells expected to express the protein that binds to the protein of the present invention (such as lung, small intestine, and heart cells of mammals), expressing the cDNAs on LB-agarose, fixing the expressed proteins onto a filter, preparing the purified protein of the present invention as a biotin-labeled or a fusion protein with the GST protein, and reacting this protein with the above filter. The desired compounds could then be detected by west western blotting using streptavidin or an anti-GST antibody (Skolnik, E. Y., Margolis, B., Mohammadi, M., Lowenstein, E., Fischer, R., Drepps, A., Ullrich, A., and Schlessinger, J. (1991) Cloning of P13 kinase-associated p85 utilizing a novel method for expression/cloning of target proteins for receptor tyrosine kinases, Cell 65: 83-90). Another method comprises the following steps. First, express the protein of the present invention fused with the SRF binding domain or the GAL4 binding. domain in yeast cells. Second, prepare a cDNA library which expresses cDNAs fused with the transcription activation domain of VP16 or GAL4 from the cells expected to express a protein that binds to the protein of the present invention. Third, introduce the cDNA into the above yeast cells. Fourth, isolate the library-derived cDNA from the positive clones. Finally, introduce the isolated cDNA into E. coli to allow it to be expressed. (When a protein that binds to the protein of the present invention is expressed in yeast cells, the reporter gene is activated and the positive clone can be detected.) This method can be performed using the two-hybrid system (MATCHMAKER Two-Hybrid system, Mammalian MATCH-MAKER Two-Hybrid Assay Kit, or MATCHMAKER One-Hybrid System (all by Clontech) orthe HybriZAP Two-Hybrid Vector System (Stratagene) (Dalton, S. and Treisman, R. (1992) Characterization of SAP-1, a protein recruited by serum response factor to the c-fos serum response element, Cell 68: 597-612). Alternatively, the binding proteins can be screened by preparing a cDNA library from the cells expected to express a substance, such as a receptor, which binds to the protein of the present invention (for example, vascular endothelial cells, bone marrow cells, or lymph duct cells), introducing it into such cells as COS, detecting the binding of the protein of the present invention by itself or labeled with a radioisotope or a fluorescence, and cloning proteins that bind to the protein of the present invention (Yamasaki, K., Taga, T., Hirata, Y., Yawata, H., Kawanishi, Y., Seed, B., Taniguchi, T., Hirano, T., and Kishimoto, T. (1988) Cloning and expression of human interleukin-6 (BSF-2/IFN beta2) receptor, Science 241: 825-828, Fukunaga, R., Ishizaka-Ikeda, E., Seto, Y., and Nagata, S. (1990) Expression cloning of a receptor for murine granulocyte colony-stimulating factor, Cell 61: 341-350). Still another method comprises applying the culture supernatant or the cellular extract of the cells expected to express a protein that binds to the protein of the present invention onto an affinity column to which the protein of the present invention has been immobilized, and purifying the proteins specifically bound to the column. In addition, a DNA encoding the protein that binds to the protein of the present invention can be obtained by determining the amino acid sequence of the binding protein, synthesizing oligonucleotides based on the sequence, and screening a cDNA library with the oligonucleotides as probes.

[0021] Furthermore, compounds that bind to the protein of the present invention can be screened by contacting compounds, a natural substance bank, or a random phage peptide display library with the immobilized protein of the present invention and detecting the molecules bound to the protein. These compounds can also be screened by high throughput screening utilizing combinatorial chemistry technology (Wrighton, N. C., Farrell, F. X., Chang, R., Kashyap, A. K., Barbone, F. P., Mulcahy, L. S., Johnson, D. L., Barrett, R. W., Jolliffe, L. K., and Dower, W. J., Small peptides as potent mimetics of the protein hormone erythropoietin, Science (United States) Jul 26 1996, 273: 458-464, Verdine, G.L., The combinatorial chemistry of nature, Nature (England) Nov 7 1996, 384: 11-13, Hogan, J.C. Jr. Directed combinatorial chemistry, Nature (England) Nov 7 1996, 384: 17-19).

[0022] VEGF-D of the present invention may be used for gene therapy by introducing the VEGF-D gene into the body of the patient with the VEGF-D deficiency, or expressing the gene in the body. An anti-sense DNA of the VEGF-D gene may also be used to inhibit the expression of the gene itself, thereby suppressing the pathological neovascularization.

[0023] Among the many available methods to introduce the VEGF-D gene or its antisense DNA into the body, the retrovirus method, the liposome method, the cationic liposome method, and the adenovirus method are preferable.

[0024] In order to express these genes in the body, the genes can be incorporated into a suitable vector and introduced into the body by the retrovirus method, the liposome method, the cationic liposome method, or the adenovirus method. Although the vectors to be used are not particularly limited, such vectors as pAdexlcw and pZIPneo are preferable.

[0025] The present invention may also be applied for diagnosing disorders caused by abnormalities of the VEGF-D gene, for example, by PCR to detect an abnormality of the nucleotide sequence of the VEGF-D gene.
[0026] Furthermore, according to the present invention, the VEGF-D protein or its agonists can be used to heal wounds, promote collateral vessel formation, and aid hematopoiesis by the hematopoietic stem cells, by taking advantage of the angiogenic effect of the VEGF-D protein. The antibodies against the VEGF-D protein or its antagonists can be used as the therapeutic agents for pathological neovascularization, lymphatic dysplasia, dyshematopoiesis, or edemas arising from various causes. The anti-VEGF-D antibodies can be used for diagnosing diseases resulting from abnormal production of VEGF-D by quantifying VEGF-D.

Brief Description of the Drawings

[0027]

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Figure 1 shows the relationship among the VEGF-D gene, the EST sequences, and the primers used for cloning. Figure 2 compares the amino acid sequences of EST (H24828) and VEGF-C.

Figure 3 compares the amino acid sequences deduced from the VEGF-D gene and from the known genes of the VEGF family proteins.

Figure 4a shows the hydrophobicity plot of VEGF-D. Figure 4b shows the prediction of the cleavage site of the VEGF-D signal peptide.

25 Best Mode for Implementing the Invention

[0028] The following examples illustrate the present invention in detail, but are not to be construed to limit the scope of the invention.

30 Example 1. Homology search by TFASTA method

[0029] The sequence CGPNKELDENTCQCVC (SEQ ID NO. 3) was designed based on the consensus sequence found in the BR3P (Balbiani ring 3 protein) repeat at the C-terminus of VEGF-C. The entire ESTs and STS sequences in the Genbank database (as of 29 February 1996) were then searched by the TFASTA method (Person and Lipman, Proc. Natl. Acad. Sci. USA 85: 2444-2448 (1988)). The searching conditions used are shown below (Table 1).

Table 1

	Sequences	392,210
	Symbols	135,585,305
	Word Size	2
	Gap creation penalty	12.0
Į	Gap extension penalty	4.0

[0030] As a result, an EST (Accession No. H24828) that is considered to code the consensus sequence was found. The sequence is one of the ESTs registered by The WashU-Merck EST Project, and nine out of 16 amino acid residues were identical. Further searching for UniGene by NCBI based on this sequence revealed that five registered sequences (T64149, H24780, H24633, H24828, and T64277 (as of 1 March 1996)), including the above EST, were considered to be derived from the same gene. T64277 and T64149, as well as H24828 and H24780, are the combination of the 5' sequence and the 3' sequence of the same dones, and the length of the insert in both of these clones was 0.9 kb (Fig. 1).

[0031] Translating the H24828 sequence into a protein sequence in a frame where homology is found suggested that this sequence codes 104 C-terminal amino acid residues. Comparing this amino acid sequence with the C-terminus of VEGF-C, 28 out of 104 amino acids (27%) were identical. Moreover, the amino acids that are important for maintaining the protein structure, such as cysteine and proline, were well conserved (Fig. 2). Conserved sequences are shown in a

black box.

Example 2. cDNA doning from a library

- Frimers for 5 RACE and 3' RACE (5' RACE primer: 5'-AGGGATGGGGAACTTGGAACGCTGAAT-3' (SEQ ID NO. 4), 3' RACE primer: 5'-GATCTAATCCAGCACCCCAAAAACTGC-3'(SEQ ID NO. 5)) were designed (Fig. 1). A double-stranded cDNA was synthesized from human lung-derived polyA* RNA using reverse transcriptase. PCR was then performed using Marathon-Ready cDNA, Lung (Chlontech), having an adapter cDNA ligated to both ends as a template cDNA, and using the above primer and adapter primer (AP-1 primer) as primers. The above adapter cDNA contains the regions to which the adapter primers AP-1 and AP-2 hybridize. The PCR was performed in a manner such that the system was exposed to treatment at 94°C for 1 min; five cycles of treatment at 94°C for 30 sec and at 72°C for 4 min; five cycles of treatment at 94°C for 20 sec and at 68°C for 4 min. (TaKaRa Ex Taq (Takara Shuzo) and the attached buffer were used as Taq polymerase instead of Advantage KlenTaq Polymerase Mix.) As a result, 1.5kb fragments were amplified at the 5' region and 0.9kb fragments at the 3' region. These fragments were cloned with the pCR-Direct Cloning System (Clontech), CR-TRAP Cloning System (Gen-Hunter), and PT7Blue-T vector (Novagen). When the 5'-RACE fragment was cloned into the pCR-Direct vector, the fragment was amplified again using 5'-CTGGTTCGGCCCAGAACTTGGAACGCTGAATCA-3' (SEQ No. 7) and 5'-CTCGCTCGCCCACTAATACGACTCACTATAGG-3' (SEQ ID NO. 8) as primers.
- 20 Example 3. Nucleotide sequence analysis

[0033] ABI PRISM Dye Terminator Cycle Sequencing Ready Reaction Kit with Amplitaq DNA Polymerase FS and 377 A DNA Sequencer (ABI) were used for DNA sequencing. The primers used are the primers in the vectors (5'-AATTAAC-CCTCACTAAAGGG-3' (SEQ ID NO. 9), 5'-CCAGGGTTTTCCCAGTCACGAC-3'(SEQ ID NO. 10)), AP-2 primer (5'-ACTCACTATAGGGCTCGAGCGGC-3' (SEQ ID NO. 11)), and 10 primers in the sequence shown below (Table 2).

Table 2

<u>·</u>	lable 2
SQ1 (SEQ ID NO. 12)	5'-AAGTCTGGAGACCTGCT-3'
SQ2 (SEQ ID NO. 13)	5'-CAGCAGGTCTCCAGACT-3'
SQ3 (SEQ ID NO. 14)	5'-CGCACCCAAGGAATGGA-3'
SQ4 (SEQ ID NO. 15)	5'-TGACACCTGGCCATTCCA-3'
SQ5 (SEQ ID NO. 16)	5'-CATCAGATGGTAGTTCAT-3'
SQ6 (SEQ ID NO. 17)	5'-ATGCTGAGCGAGAGTCCATA-3'
SQ7 (SEQ ID NO. 18)	5'-CACTAGGTTTGCGGCAACTT-3'
SQ8 (SEQ ID NO. 19)	5'-GCTGTTGGCAAGCACTTACA-3'
SQ9 (SEQ ID NO. 20)	5'-GATCCATCCAGATCCCTGAA-3'
SQ10 (SEQ ID NO. 21)	5'-CAGATCAGGGCTGCTTCTA-3'
	SQ2 (SEQ ID NO. 13) SQ3 (SEQ ID NO. 14) SQ4 (SEQ ID NO. 15) SQ5 (SEQ ID NO. 16) SQ6 (SEQ ID NO. 17) SQ7 (SEQ ID NO. 18) SQ8 (SEQ ID NO. 19) SQ9 (SEQ ID NO. 20)

[0034] Determining the nucleotide sequence of the 1.5kb fragment at the 5'-side and the 0.9kb fragment at the 3'-side revealed that the sequence of the overlapping region was identical, confirming that 5'- and 3'-side cDNAs of the desired gene were obtained. Determining the entire nucleotide sequence of the cDNA revealed that this novel gene has the full length of 2 kb and can code a protein consisting of 354 amino acid residues (SEQ ID NO. 1 and SEQ ID NO. 2). Figure 1 shows the relation between this gene and the EST sequences registered in the Genbank database. Comparing the amino acid sequence with other VEGF family proteins revealed that the amino acids that are well conserved between family proteins are also conserved in this novel gene, and therefore this gene is obviously a new member of the VEGF family (Fig. 3). In Fig. 3, HSVEGF indicates human VEGF; HSVEGF-D, HSVEGF-C, and HSVEGF-B indicate human VEGF homologues (human VEGF-D, human VEGF-C, and human YEGF-B, respectively); HSPDGF-A indicates human PDGF-A; HSPDGF-B indicates human PDGF-B; and HSP1GF2 indicates human P1GF2. The conserved sequences are shown in a black box. Since VEGF-D is highly homologous to VEGF-C that was cloned as the Flt4 ligand, it was presumed to be a ligand to a Flt-4-like receptor.

[0035] Deducing the signal peptide cleavage site (Fig. 4b) by hydrophobicity plot (Fig. 4a) and the method of von Heijne (von Heijne, G, Nucleic Acids Res. 14, 4683-4690(1986)), N-terminal 21 amino acid residues may be cleaved as

signal peptides, and they may also undergo additional processing like VEGF-C.

Example 4. Northern blot analysis

[0036] A 1kb fragment, which had been cut out by digestion with EcoRI from the 5'-fragment subcloned into pCR-Direct vector, was labeled with[α-32P]dCTP and used as a probe. Labeling was performed by random priming using Ready-to Go DNA labeling beads (Pharmacia). Hybridization was performed in ExpressHyb Hybridization Solution (Clontech) by the usual method using Multiple Tissue Northern (MTN) Blot-Human, Human II, Human Fetal, and Human Cell lines (Clontech). Significant expression was observed in lung, heart, and intestine. Weak expression was observed in skeletal muscle, ovary, colon, and pancreas. The apparent molecular weight of the mRNA was 2.2 kb, and the cloned fragment seemed to be almost the full length of the gene.

Example 5. VEGF-D protein expression in E. coli

15 [0037] Two primers, 5'-TCCAGATCTTTTGCGGCAACTTTCTATGACAT-3' (SEQ ID NO. 22) and 5'-CAGGTCGACT-CAAACAGGCACTAATTCAGGTAC-3' (SEQ ID NO. 23), were synthesized to amplify the region corresponding to the 89th to 181st amino acid residues of human VEGF cDNA. The thus-obtained DNA fragment was digested with restriction enzymes BgllI and Sall, and ligated using ligation kit II (Takara Shuzo Co., Ltd) to plasmid pQE42 ((QIAGEN), which had been digested with restriction enzymes BamHI and Sall. The resulting plasmid was introduced into E. coli SG19003[pREP4] (QIAGEN), and a plasmid, which was obtained as designed without any mutation, was selected (pQE42-BS3). Plasmid pQE42-BS3 was introduced into E. coli BL21 (Invitorogen) and cultured in 10 ml of L Broth containing 100 mg/l bicucilline (ampicillin sodium for injection, Meiji Seika Kaisha, Ltd.). 200 ml of fresh L Broth was then inoculated with the culture. After incubation at 37°C for 1.5 hours, IPTG was added to 3 mM, and the culture was further incubated at 37°C for 5 hours. After cells were harvested, a protein was purified with a Ni-NTA column following the protocol of QIAexpress Typell kit.

Example 6. Expression of DHFR-VEGF-D fusion protein in E. coli

[0038] The region corresponding to the 89th to 181st amino acid residues of human VEGF cDNA was amplified with the same primers used in Example 5. The thus-obtained DNA fragment was digested with restriction enzymes Bgll and Sall. The fragment was then ligated using ligation kit II (Takara Shuzo Co., Ltd.) to the plasmid pQE40 (QIAGEN), which had been digested with restriction enzymes BamHI and Sall. The resulting plasmid was introduced into E. coli SG19003[pREP4] (QIAGEN), and a plasmid, which was obtained as designed without any mutation, was selected (pQE40-BS3). Plasmid pQE40-BS3 was introduced into E. coli BL21 (Invitrogen) and cultured in 10 ml of L Broth containing 100 mg/l bicucilline (ampicillin sodium for injection, Meiji Seika Kaisha, Ltd.). 200 ml of fresh L Broth was then inoculated with the culture. After incubation at 37°C for 1.5 hours, IPTG was added to 3mM, and the culture was further incubated at 37°C for 5 hours. After cells were harvested, a DHFR-VEGF-D fusion protein was purified with a Ni-NTA column following the protocol of a QIAexpress TypeII kit.

40 Example 7. Cloning mouse VEGF-D cDNA

[0039] Two Hybond-N+ (Amersham) filters (20 cm x 22 cm) on which 1.5 x 10⁵ pfu of Mouse lung 5'-stretch cDNA library was transferred were prepared. Gradient hybridization from 68°C to 55°C was performed for 2 hours in ExpressHyb Hybridization Solution (Clontech) using as a probe an approximately 50 ng Pvu II fragment of human VEGF-D, which had been labeled with α³²P-dCTP (Amersham) using Ready-To-Go DNA Labeling Beads(-dCTP) (Pharmacia). The filters were washed four times in 2 x SSC, 0.05% SDS at room temperature for 10 min, then washed in 0.1 x SSC, 0.1% SDS at 45°C for 3 min. The washed filters were exposed overnight at -80°C using HyperFilm MP (Amersham) and intensifying paper. Positive clones were subjected to the second screening in the same manner as above to isolate a single clone. Isolated lambda DNAs were purified from the plate lysate using a QIAGEN Lambda MAX I Kit (Qiagen). Insert DNAs were cut out with EcoRI and subcloned into pUC118 EcoRI/BAP (Takara Shuzo Co., Ltd.). Its nucleotide sequence was then determined with ABI377 sequencer (Perkin Elmer). The cDNA coding the full length of mouse VRGF-D was reconstructed with two of the obtained clones that overlapped each other. SEQ ID NO. 24 shows the nucleotide sequence of mouse VEGF-D cDNA and the deduced amino acid sequence therefrom.

55 Example 8. Cloning rat VEGF-D cDNA

[0040] Two Hybond-N+ (Amersham) filters (20 cm x 22 cm), on which 1.5 x 10⁵ pfu of Rat lung 5'-stretch cDNA library had been transferred, were prepared. Gradient hybridization from 68°C to 55°C was performed for 2 hours in

ExpressH.Fyb Hybridization Solution (Clontech) using as a probe an approximately 1 μg fragment containing 1-782 bp of the mouse VEGF-D cDNA which had been labeled with α³²P-dCTP (Amersham) using Ready-To-Go DNA Labeling Beads(-dCTP) (Pharmacia). The filters were washed four times in 2 x SSC, 0.05% SDS at room temperature for 10 min, then washed in 0.1 x SSC, 0.1% SDS at 45°C for 3 min. The washed filters were exposed overnight at -80°C using HyperFilm MP (Amersham) and intensifying paper. Positive clones were subjected to the second screening in the same manner as above to isolate a single clone. The isolated positive clone was excised into pBluescript using E. coli SOLAR (Stratagene) and helper phage ExAssist (Stratagene), then the sequence was determined with ABI377 sequencer (Perkin Elmer). The sequence seemed to be the rat VEGF-D cDNA but did not contain the termination codon.

[0041] To obtain the C-terminal cDNA which had not been obtained, PCR was performed using Marathon-Ready rat kidney cDNA (Clontech) as a template and 5' primerGCTGCGAGTGTCTGTAAA (SEQ ID NO. 26) and 3' primer GGGTAGTGGGCAACAGTGACAGCAA (SEQ ID NO. 27) with 40 cycles of 94°C for 15 sec, 55°C for 30 sec, and 72 °C for 2 min. After the thus-obtained fragment was subcloned into pGEM-T vector (promega), the nucleotide sequence was determined with ABI377 sequencer (Perkin Elmer). The resulting clone contained the C-terminus of rat VEGF-D. Based on the results of sequencing the clone obtained by plaque hybridization and the clone obtained by PCR, the full length of the rat VEGF-D sequence was determined. SEQ ID NO. 25 shows the determined nucleotide sequence and the deduced amino acid sequence therefrom.

Industrial Applicability

[0042] In the present invention, a novel protein (VEGF-D) having significant homology to VEGF-C and its gene have been isolated. VEGF-D appears to be involved in the pathological neovascularization associated with diabetes, rheumatoid arthritis, the growth of solid tumors, differentiation and proliferation of blood cells, formation of lymphatic vessels, and formation of edema resulting from various causes as well as the normal neovascularization at the developmental stage. The gene of the present invention can be used to diagnos disorders caused by abnormalities of the VEGF-D gene and gene therapy for the VEGF-D deficiency. The VEGF-D protein, which is obtained by expressing the gene of the present invention, can be used for healing wounds, promoting collateral vessel formation, and aiding hematopoietic stemcell proliferation. The antibodies or inhibitors against the VEGF-D protein can be used for treating angiodysplasia and lymphangiodysplasia associated with inflammation, edemas arising from various causes, dyshematopoiesis, and, as a novel anticancer agent, for treating pathological neovascularization. The VEGF-D protein and its antibodies can be useful for diagnosing diseases resulting from abnormal production of VEGF-D.

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Sequence Listing

(1) Name or appellation of Applicant: Chugai Research Institute for 5 Molecular Medicine, Inc. (2) Title of the Invention: Novel VEGF-like Factor (3) Reference Number: C1-802PCT 10 (4) Application Number: (5) Filing date: (6) Country where the priority application was filed and the application number of the application: Japan, No. Hei 8-185216 15 (7) Priority date: July 15, 1996 (8) Number of sequences: 27 20 SEQ ID NO: 1 SEQUENCE LENGTH: 354 SEQUENCE TYPE: amino acid TOPOLOGY: linear 25 MOLECULE TYPE: protein ORIGINAL SOURCE: ORGANISM: Homo sapiens 30 TISSUE TYPE: lung SEQUENCE DESCRIPTION: Met Tyr Arg Glu Trp Val Val Val Asn Val Phe Met Met Leu Tyr Val 5 10 15 35 Gln Leu Val Gln Gly Ser Ser Asn Glu His Gly Pro Val Lys Arg Ser 20 25 30 Ser Gln Ser Thr Leu Glu Arg Ser Glu Gln Gln Ile Arg Ala Ala Ser 40 35 45 Ser Leu Glu Glu Leu Leu Arg Ile Thr His Ser Glu Asp Trp Lys Leu 55 Trp Arg Cys Arg Leu Arg Leu Lys Ser Phe Thr Ser Met Asp Ser Arg 70 75 80 65 Ser Ala Ser His Arg Ser Thr Arg Phe Ala Ala Thr Phe Tyr Asp Ile . 90 85

Glu Thr Leu Lys Val Ile Asp Glu Glu Trp Gln Arg Thr Gln Cys Ser

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100

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•	Pro	Arg	Glu	Thr	Cys	Val	Glu	Val	Ala	Ser	Glu	Leu	Gly	Lys	Ser	Thr
			115					120					125	5	•	
5	Asn	Thr	Phe	Phe	Lys	Pro	Pro	Cys	Val	Asn	Val	Phe	Arg	Cys	Gly	Gly
		130					135	5				140				
	Cys	Cys	Asn	Glu	Glu	Ser	Leu	Ile	Cys	Met	Asn	Thr	Ser	Thr	Ser	Tyr
10	145					·150)				155			٠		160
	Ile	Ser	Lys	Gln	Leu	Phe	Glu	Ile	Ser	Val	Pro	Leu	Thr	Ser	Val	Pro
					165					170					175	5
	Glu	Leu	Val	Pro	Val	Lys	Val	Ala	Asn	His	Thr	Gly	Cys	Lys	Cys	Leu
15				180					185					190	١	
	Pro	Thr	Ala	Pro	Arg	His	Pro	Tyr	Ser	Ile	Ile	Arg	Arg	Ser	Ile	Gln
			195					200					205			
20	Ile	Pro	Glu	Glu	Asp	Arg	Cys	Ser	His	Ser	Lys	ГÀЗ	Leu	Cys	Pro	Ile
		210					215					220				
		Met	Leu	Trp	Asp	Ser	Asn	Lys	Сув	Lys	Суз	Val	Leu	Gln	Glu	Glu
	225					230					235					240
25	Asn	Pro	Leu	Ala		Thr	Glu	Asp	His	Ser	His	Leu	Gln	Glu	Pro	Ala
			_		245					250					255	
	Leu	Cys		Pro	His	Met	Met	Phe		Glu	Asp	Arg	Cys	Glu	Cys	Val
30		_		260		_	_		265	_				270		
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TOPOLOGY: linear

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	CCT	GCGG	CAT	ACAT'	TGGA(GA G	ATTT	T TT T	TA A	TTTC'	TGGA	CAY	GAAG	TAA	ATTT	AGAGTG	120
o	CTT	TCYA	ATT '	TCAG	GTAG.	AA G	ACAT	GTCC	A CC	rtct(GATT	ATT	TTTG	GAG	AACA	TTTTGA	180
	TTT	TTTT	CAT	CTCT	CTCT	cc c	CACC	CCTA	A GA	rtgt(GCAA	AAA	AAGC	GTA	CCTT	GCCTAA	240
	TTG	AAAT.	TAA	TTCA	TTGG.	AT T	TTGA'	TCAG	A AC	rgat(CATT	TGG	TTTT	CTG	TGTG.	AAGTTT	300
	TGA	GGTT'	TCA .	AACT'	TTÇC	IT C	TGGA	GAAT	G CC	r tt t(GAAA	CAA	TTTT	CTC	TAGC'	TGCCTG	360
5	ATG	TCAA	CTG	CTTA	GTAA'	TC A	GTGG	ATAT:	r ga	AATA'	TCA	AA .	ATG '	TAC	AGA (GAG	414
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o	TGG	GTA	GTG	GTG	aat	GTT	TTC	ATG	ATG	TTG	TAC	GTC	CAG	CTG	GTG	CAG	462
	Trp	Val	Val	Val	Asn	Val	Phe	Met	Met	Leu	Tyr	Val	Gln	Leu	Val	Gln	
	5					10 .					15					20	
	GGC	TCC	AGT	AAT	GAA	CAT	GGA	CCA	GTG	AAG	CGA	TCA	TCT	CAG	TCC	ACA	510
5	Gly	Ser	Ser	Asn	Glu	His	Gly	Pro	Val	Lys	Arg	Ser	Ser	Gln	Ser	Thr	
					25					30					35		
	TTG	GAA	CGA	TCT	GAA	CAG	CAG	ATC	AGG	GCT	GCT	TCT	AGT	TTG	GAG	GAA	558
o	Leu	Glu	Arg	Ser	Glu	Gln	Gln	Ile	Arg	Ala	Ala	Ser	Ser	Leu	Glu	Glu	
-				40					45					50			
	CTA	CTT	CGA	ATT	ACT	CAC	TCT	GAG	GAC	TGG	AAG	CTG	TGG	AGA	TGC	AGG	606
	Leu	Leu	Arg	Ile	Thr	His	Ser	Glu	Asp	Trp	Lys	Leu	Trp	Arg	Cys	Arg	
5			55					60					65				
	CTG	AGG	CTC	AAA	AGT	TTT	ACC	AGT	ATG	GAC	TCT	CGC	TCA	GCA	TCC	CAT	654
	Leu	Arg	Leu	Lys	Ser	Phe	Thr	Ser	Met	qzA	Ser	Arg	Ser	Ala	Ser	His	
0		70					75					80					
-	CGG	TCC	ACT	AGG	TTT	GCG	GCA	ACT	TTC	TAT	GAC	ATT	GAA	ACA	CTA	AAA	702
	Arg	Ser	Thr	Arg	Phe	Ala	Ala	Thr	Phe	Tyr	qzA	Ile	Glu	Thr	Leu	Lys	

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	GTT	ATA	GAT	GAA	GAA	TGG	CAA	AGA	ACT	CAG	TGC	AGÇ	сст	AGA	GAA	ACG	750
5	Val	Ile	Asp	Glu	Glu	Trp	Gln	Arg	Thr	Gln	Cys	Ser	Pro	Arg	Glu	Thr	
					105	5				110)				115		
	TGC	GTG	GAG	GTG	GCC	AGT	GAG	CTG	GGG	AAG	AGT	ACC	AAC	ACA	TTC	TTC	798
10	Суз	Val	Glu	Val	Ala	Ser	Glu	Leu	Gly	Lys	Ser	Thr	Asn	Thr	Phe	Phe	
				120					125					130	0		
																GAA	846
	Lys	Pro	Pro	Сλа	Val	Asn	Val	Phe	Arg	Суѕ	Gly	Gly	Сла	Cys	Asn	Glu	
15			135					140					145				
																CAG	894
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20		150					155					160					
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			215					220					225			•	
	GAT	AGC	AAC	AAA	TGT	AAA	TGT	GTT	TTG	CAG	GAG	GAA	AAT	CCA	СТТ	GCT	1134
40	Asp	Ser	Asn	Lys	Cys	Lys	Суз	Val	Leu	Gln	Glu	Glu	Asn	Pro	Leu	Ala	
10		230					235					240					
	GGA	ACA	GAA	GAC	CAC	TCT	CAT	CTC	CAG	GAA	CCA	GCT	CTC	TGT	GGG	CCA	1182
	Gly	Thr	Glu	Asp	His	Ser	His	Leu	Gln	Glu	Pro	Ala	Leu	Cys	Gly	Pro	
15 ·	245					250					255					260	
	CAC	ATG	ATG	TTT	GAC	GAA	GAT	CGT	TGC	GAG	TGT	GTC	TGT	AAA	ACA	CCA	1230
	His	Met	Met	Phe	Asp	Glu	Asp	Arg	Cys	Glu	Cys	Val	Суз	Lys	Thr	Pro	
50					265					270					275		
	TGT	CCC	AAA	GAT	CTA	ATC	CAG	CAC	ccc	AAA.	AAC	TGC	agt	TGC	TTT	GAG	1278
	Cys	Pro	Lys	Asp	Leù	Ile	Gln	His	Pro	Lys	Asn	Cys	Ser	Cys	Phe	Glu	

	280 285 290	
	TGC AAA GAA AGT CTG GAG ACC TGC TGC CAG AAG CAC AAG CTA TTT CAC	1326
5	Cys Lys Glu Ser Leu Glu Thr Cys Cys Gln Lys His Lys Leu Phe His	
	295 300 305	
	CCA GAC ACC TGC AGC TGT GAG GAC AGA TGC CCC TTT CAT ACC AGA CCA	1374
10	Pro Asp Thr Cys Ser Cys Glu Asp Arg Cys Pro Phe His Thr Arg Pro	
	310 315 320	
	TGT GCA AGT GGC AAA ACA GCA TGT GCA AAG CAT TGC CGC TTT CCA AAG	1422
15	Cys Ala Ser Gly Lys Thr Ala Cys Ala Lys His Cys Arg Phe Pro Lys	
	325 330 335 340	
	GAG AAA AGG GCT GCC CAG GGG CCC CAC AGC CGA AAG AAT CCT	1464
	Glu Lys Arg Ala Ala Gln Gly Pro His Ser Arg Lys Asn Pro	
20	345 350	
	TGATTCAGCG TTCCAAGTTC CCCATCCCTG TCATTTTTAA CAGCATGCTG CTTTGCCAAG	1524
	TTGCTGTCAC TGTTTTTTC CCAGGTGTTA AAAAAAAAA CCATTTTACA CAGCACCACA	1584
25	GTGAATCCAG ACCAACCTTC CATTCACACC AGCTAAGGAG TCCCTGGTTC ATTGATGGAT	1644
	GTCTTCTAGC TGCAGATGCC TCTGCGCACC AAGGAATGGA GAGGAGGGGA CCCATGTAAT	1704
	CCTTTTGTTT AGTTTTGTTT TTGTTTTTTG GTGAATGAGA AAGGTGTGCT GGTCATGGAA	1764
30	TGGCAGGTGT CATATGACTG ATTACTCAGA GCAGATGAGG AAAACTGTAG TCTCTGAGTC CTTTGCTAAT CGCAACTCTT GTGAATTATT CTGATTCTT TTTATGCAGA ATTTGATTCG	1824
	TATGATCAGT ACTGACTTTC TGATTACTGT CCAGCTTATA GTCTTCCAGT TTAATGAACT	1884
	ACCATCTGAT GTTTCATATT TAAGTGTATT TAAAGAAAAT AAACACCATT ATTCAAGTCT	1944
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35	SEQ ID NO: 3	
	SEQUENCE LENGTH: 16	
	SEQUENCE TYPE: amino acid	
40	TOPOLOGY: linear	
	MOLECULE TYPE: peptide	
	ORIGINAL SOURCE:	
4 5	ORGANISM: Homo sapiens	
	TISSUE TYPE: lung	
	SEQUENCE DESCRIPTION:	•
	Cys Gly Pro Asn Lys Glu Leu Asp Glu Asn Thr Cys Gln Cys Val Cys	
50	1 5 10 15	

	SEQ ID NO: 4	
	SEQUENCE LENGTH: 27	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: other-nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	AGGGATGGGG AACTTGGAAC GCTGAAT	27
15	SEQ ID NO: 5	٠
	SEQUENCE LENGTH: 27	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
25	GATCTAATCC AGCACCCCAA AAACTGC	27
	SEQ ID NO: 6	
30	SEQUENCE LENGTH: 27	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
<i>35</i>	TOPOLOGY: linear	
00	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CCATCCTAAT ACGACTCACT ATAGGGC	27
40		·
	SEQ ID NO: 7	
	SEQUENCE LENGTH: 33	
45	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
50	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	•
	CTGGTTCGGC CCAGAACTTG GAACGCTGAA TCA	33

	SEQ ID NO: 8	
	SEQUENCE LENGTH: 32	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CTCGCTCGCC CACTAATACG ACTCACTATA GG	32
15		
	SEQ ID NO: 9	
	SEQUENCE LENGTH: 20	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
25	SEQUENCE DESCRIPTION:	
	AATTAACCCT CACTAAAGGG	20
	SEQ ID NO: 10	
30	SEQUENCE LENGTH: 22	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
35	TOPOLOGY: linear	
•	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	•
	CCAGGGTTTT CCCAGTCACG AC	22
40	·	
	SEQ ID NO: 11	
	SEQUENCE LENGTH: 23	
45	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
50	SEQUENCE DESCRIPTION:	
	ACTCACTATA GGGCTCGAGC GGC	23

	SEQ ID NO: 12	
	SEQUENCE LENGTH: 17	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	AAGTCTGGAG ACCTGCT	17
15		
	SEQ ID NO: 13	
	SEQUENCE LENGTH: 17	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
25	SEQUENCE DESCRIPTION:	
	CAGCAGGTCT CCAGACT	17
	SEQ ID NO: 14	
30	SEQUENCE LENGTH: 17	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
35	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CGCACCCAAG GAATGGA	17
10		
	SEQ ID NO: 15	
	SEQUENCE LENGTH: 18	
15	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
0	SEQUENCE DESCRIPTION:	
	TGACACCTGG CCATTCCA	18

	SEQ ID NO: 16	
	SEQUENCE LENGTH: 18	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CATCAGATGG TAGTTCAT	18
45		
15	SEQ ID NO: 17	
	SEQUENCE LENGTH: 20	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
<i>2</i> 5	SEQUENCE DESCRIPTION:	
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	SEQ ID NO: 18	
30	SEQUENCE LENGTH: 20	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
35	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
40	CACTAGGTTT GCGGCAACTT	20
	SEQ ID NO: 19	
	SEQUENCE LENGTH: 20	
45	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
50	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	GCTGTTGGCA AGCACTTACA	20

	SEQ ID NO: 20	
	SEQUENCE LENGTH: 20	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	GATCCATCCA GATCCCTGAA	20
15		
	SEQ ID NO: 21	
	SEQUENCE LENGTH: 19	
•	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
25	SEQUENCE DESCRIPTION:	
	CAGATCAGGG CTGCTTCTA	19
	SEQ ID NO: 22	
30	SEQUENCE LENGTH: 32	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
35	TOPOLOGY: linear	
33	MOLECULE TYPE: other nucleic acid, synthetic DNA	
	SEQUENCE DESCRIPTION:	
	TCCAGATCTT TTGCGGCAAC TTTCTATGAC AT	32
10		
	SEQ ID NO: 23	
	SEQUENCE LENGTH: 33	
4 5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: other nucleic acid, synthetic DNA	
50	SEQUENCE DESCRIPTION:	
	CAGGTCGACT CAAACAGGCA CTAATTCAGG TAC	33

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25										1	Met 1	ryr (Gly	Glu '	Trp	Gly	
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												TTG					161
<i>30</i>	Met	Gly	Asn		Leu	Met	Met	Phe		Val	Tyr	Leu	Val		Gly	Phe	
	-00			10					15					20			
						•										TCC	209
	Arg	zet	25	HIS	GIY	Pro	Vai		Asp	hue	Ser	Phe		Arg	Ser	Ser	
35	ccc	ጥርሮ		ጥጥር	CAA	CCB	ጥርጥ	30	C	CRC	3 M.C	CC >	35	ccm	mca	100	262
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	Arg	, Glu	Thi	Cys	Val	Glu	Val	Ala	Ser	Glu	Lev	ı Gly	Lys	Th	r Th	r Asn	
10		120				•	125					130					
																C TGC	
	Thr	Phe	Phe	Lys	Pro	Pro	Cys	Val	Asn	Val	Phe	arg	Суз	Gly	y Gly	y Cys	
	135					140					145					150	
15																ATC	
	Cys	Asn	Glu	Glu	Gly	Val	Met	Суз	Met	Asn	Thr	Ser	Thr	Sez	ту	lle	
					155					160					165		
20																GAG	641
	Ser	Lys	Gln		Phe	Glu	Ile				Leu	Thr	Ser	Val	Pro	Glu	
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																ccc	689
25	Leu	vai			Lys	IIe	Ala		His	Thr	Gly			Cys	Leu	Pro	
	ACG	ccc	185		Cam	ccm	m > 4	190					195				
												AGA					737
30	****	200		nry	птэ	PLO	205	ser	116	TTE	Arg	Arg	Ser	Ile	Gln	Thr	
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40	CCA	CTG	ССТ	GGG	ACA	GAA	GAC	CAC	TCT		CTC	CAG	GAA	רכר		CTC	001
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				250			•		255	-1-			724	260		neu	
45	TGT	GGA	CCG	CAC	ATG	ACG	TTT	GAT		GAT	CGC	TGT :	GAG			ጥርም	0.20
7 5												Cys					929
	-	·	265					270			3	-1-	275	-13	- al	Cys	
	AAA	GCA	CCA	TGT	CCG	GGA (ATT	CAG	CAC	CCG (_	226	ምር ሮ	acm	077
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	TGC TTT GAG TGC AAA GAA AGT CTG GAG AGC TGC TGC CAA AAG CAC AAG	1025
	Cys Phe Glu Cys Lys Glu Ser Leu Glu Ser Cys Cys Gln Lys His Lys	
5	295 300 305 310	
	ATT TTT CAC CCA GAC ACC TGC AGC TGT GAG GAC AGA TGT CCT TTT CAC	1073
	Ile Phe His Pro Asp Thr Cys Ser Cys Glu Asp Arg Cys Pro Phe His	
10	315 · 320 325	
	ACC AGA ACA TGT GCA AGT AGA AAG CCA GCC TGT GGA AAG CAC TGG CGC	1121
	Thr Arg Thr Cys Ala Ser Arg Lys Pro Ala Cys Gly Lys His Trp Arg	
	330 335 340	
15	TTT CCA AAG GAG ACA AGG GCC CAG GGA CTC TAC AGC CAG GAG AAC CCT	1169
	Phe Pro Lys Glu Thr Arg Ala Gln Gly Leu Tyr Ser Gln Glu Asn Pro	
	345 350 355	
20	TGATTCAACT TCCTTTCAAG TCCCCCCATC TCTGTCATTT TAAACAGCTC ACTGCTTTGT	1229
	CAAGTTGCTG TCACTGTTGC CCACTACCCC TGCCCCCCCC CCCCCCGCC TCCAGGTGTT	1289
·	AGAAAAGTTG ATTTGACCTA GTGTCATGGT AAAGCCACAT TTCCATGCAA TGGCGGCTAG	1349
	GTGATTCCCC AGTTCACTGA CAAATGACTT GTAGCTTCAA ATGTCTTTGC GCCATCANCA	1409
25	CTCAAAAAGG AAGGGGTCTG AAGAACCCCT TGTTTGATAA ATAAAAACAG GTGCCTGAAA	1469
	CAAAATATTA GGTGCCACTC GATTGGGTCC CTCGGGCTGG CCAAATTCCA AGGGCAATGC	1529
	TCCTGAATTT ATTGTGCCCC TTCCTTAATG CGGAATTTCC TTTTGTTTGA TT	1581
<i>30</i>	· · · · · · · · · · · · · · · · · · ·	
	SEQ ID NO: 25	
	SEQUENCE LENGTH: 1491 .	
	SEQUENCE TYPE: nucleic acid	
35	TOPOLOGY: linear	
	MOLECULE TYPE: cDNA to mRNA	
	ORIGINAL SOURCE:	
40	ORGANISM: rat	
	TISSUE TYPE: lung	
	FEATURE:	
45	NAME/KEY: CDS	
45	LOCATION: 2701247	
	IDENTIFICATION METHOD: E	
	SEQUENCE DESCRIPTION:	
50	GCCACCTCTT GATTATTTGT GCAGCGGGAA ACTTTGAAAT AGTTTTCATC TCTTTCTCCC	60
	ATACTAAGAT TGTGTGTGC CGTGGGGGAG TCCTTGACTA ACTCAAGTCA TTTCATTGGA	120
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	TT	TTGA	TAC	AACT	OTADI	AT (GTGAT	rtta?	T TI	TCCA	TGT	DAA	TTT	rggg	GCT	CAAAC	T 180
	TT	GCTT	CTGG	AGA	ATGC	TT 1	TGC	ACAC	T T	TCAG	TAGO	TGC	CTG	AAA	CAA	TGCTT.	A 240
5	GC	CATC	AGTG	GACA	ATTTC	AA A	TAT	CAAA	ATO	TAT	GGA	GAG	TG	GCC	C GC	A GTG	293
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10	AA	r att	CTC	ATC	ATG	TCC	TAT	GTG	TAC	CTG	GTG	CAG	GGC	TTC	AG2	TTA T	341
	Ası	n Ile	e Leu	ı Met	: Met	Ser	Tyr	. Val	Tyr	Leu	Val	Gln	Gly	Phe	s Ser	: Ile	
		10					15					20			•		
																TCT	389
15			Arg	y Ala	Val	Lys	Asp	Val	Ser	Leu	Glu	Arg	Set	Ser	Arg	Ser	
	25					30					35					40	
																GAA	437
20	Val	Lev	Glu	Arg		Glu	Gln	Gln	Ile	Arg	Ala	Ala	Ser	Thr	Leu	Glu	
	•				45					50					55		
																TGC	485
	GIA	Leu	Leu		Val	Ala	His	Ser		Asp	Trp	rys	Leu	_	Arg	Cys	
25		mm/s		60					65					70			
				CTT													533
	Arg	Leu	туs 75	Leu	rÀ2	Ser	Leu		Asn	Val	Asp	Ser		Ser	Thr	Ser	
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				ACC													581
		90	JGL	Thr	ALY	rne	95	WIG	THE	Pne	туг		Thr	GTA	Thr	Leu	
	AAA		АТА	GAT	GAA	GAA		CNG	ncc.	A C C	CAA	100	100				4.5.
35				Asp													629
	105			·.op	V	110	110	OIII	My	1111	115	cys	Ser	PIO	Arg		
			GTA	GAA	GTC		AGT	GAG	ርተር	GGG		ara	ACC.	220	N.C.B	120	627
40				Glu													677
		-			125					130	- ,-			AGH	135	rne	
	TTC	AAG	ccc	CCT		GTA	AAT	GTC	TTC		TGT	GGA	GGA	TGC		ልልጥ	725
45				Pro													725
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	GAA	GAG	AGC	GTG	ATG	TGT	ATG	AAC		AGC	ACC	TCC	TAC		ጥርር	אא	773
				Val													113
50			155			•	_	160				-	165			~ _J 3	
	CAG	CTC	TTT	GAG	ATA	TCA	GTG		CTG	ACA	TCA	GTG		GAG	TTA	GTG	821
																	- 2 1

	Gln	Leu	Phe	Glu	Ile	Ser	Val	Pro	Leu	Thr	Ser	Val	Pro	Glu	Leu	Val	
		170					175					180					
5	CCT	GTT	AAA	ATT	GCC	AAC	CAT	ACG	GGT	TGT	AAG	TGT	TTG	ccc	ACG	GGC	869
	Pro	Val	Lys	Ile	Ala	Asn	His	Thr	Gly	Cys	Lys	Cys	Leu	Pro	Thr	Gly	
	185					190					195					200	
10	ccc	CGG	CAT	CCT	TAT	TCA	ATT	ATC	AGA	AGA	TCC	ATT	CAG	ATC	CCA	GAA	917
	Pro	Arg	His	Pro	Tyr	Ser	Ile	Ile	Arg	Arg	Ser	Ile	Gln	Ile	Pro	Glu	
					205					210					215	i	
	GAA	GAT	CAÁ	TGT	CCT	CAT	TCC	AAG	AAA	CTC	TGT	CCT	GTT	GAC	ATG	CTG	965
15	Glu	Asp	Gln	Cys	Pro	His	Ser	Lys	Lys	Leu	Cys	Pro	Val	Asp	Met	Leu	
				220					225					230			
	TGG	GAT	AAC	ACC	AAA	TGT	AAA	TGT	GŢŢ	TTA	CAA	GAT	GAG	AAT	CCA	CTG	1013
20	Trp	Asp	Asn	Thr	Lys	Cys	Lys	Cys	Val	Leu	Gln	Asp	Glu	Asn	Pro	Leu	
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	CCT	GGG	ACA	GAA	GAC	CAC	TCT	TAC	CTC	CAG	GAA	ccc	GCT	CTC	TGT	GGA	1061
<i>25</i>	Pro	Gly	Thr	Gl u	qeA	His	Ser	Tyr	Leu	Gln	Glu	Pro	Ala	Leu	Cys	Gly	
20		250					255					260					
	CCA	CAC	ATG	ATG	TTT	GAT	GAA	GAT	CGC	TGC	GAG	TGT	GTC	TGT	AAA	GCA	1109
	Pro	His	Met	Met	Phe	qzA	Glu	Asp	Arg	Суз	Glu	Cys	Val	Cys	Lys	Ala	
30	265					270					275					280	
	CCA	TGT	CCT	GGA	GAT	CTC	ATT	CAG	CAC	CCG	GAA	AAC	TGC	AGT	TGC	TTT	1157
	Pro	Суз	Pro	Gly	Asp	Leu	.Ile	Gln	His	Pro	Glu	Asn	Cys	Ser	Cys	Phe	
35					285					290					295		
	GAA	TGC	AAA	GAA	AGT	CTG	GAA	AGC	TGT	TGC	CAA	AAG	CAC	AAG	ATG	TTT	1205
	Glu	Суз	Lys	Glu	Ser	Leu	Glu	Ser	Cys	Cys	Gln	Lys	His	Lys	Met	Phe	
40				300					305					310			
40	CAC	CCT	GAC	ACC	TGC	AGA	TCA	ATG	GTC	TTT	TCA	CTG	TCC	CCT			1247
	His	Pro	Ąsp	Thr	Cys	Arg	Ser	Met	Val	Phe	Ser	Leu	Ser	Pro			
			315					320					325				
45	TAAT	TTG	STT 1	PACTO	GTG!	AC AT	LATT	\AGG#	A CAT	ACTA	LACC	TGAT	TTAT	TTG (GGC	CTTTT	1307
	CTCI	CAG	GC (CAAC	CAC	AC TO	CTTA	\AGG!	A ACI	CAGA	CGT	TTG	CCTC	TA A	AGAAI	TACAT	1367
	GGA	GTA	TA T	ragac	TGAT	rg an	taa ti	ATTGT	CTI	CTTO	TTT	CAA	ACAGO	GT (TCA	GATTA .	1427
50	CAGA	ACCC	STA 1	rrgco	CATG	CC TO	GCCG1	CATO	CT!	ATCAT	CGAG	CGG	DAAA	AA 1	CAC	TGGCAT	1487
	TTAF	\															1491

SEQ ID NO: 26

SEQUENCE LENGTH: 20

SEQUENCE TYPE: nucleic acid

STRANDEDNESS: single

TOPOLOGY: linear

MOLECULE TYPE: other nucleic acid, synthetic DNA

SEQUENCE DESCRIPTION:

GCTGCGAGTG TGTCTGTAAA

20

15

SEQ ID NO: 27

SEQUENCE LENGTH: 25

SEQUENCE TYPE: nucleic acid

20 STRANDEDNESS: single

TOPOLOGY: linear

MOLECULE TYPE: other nucleic acid, synthetic DNA.

SEQUENCE DESCRIPTION:

GGGTAGTGGG CAACAGTGAC AGCAA

25

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Claims

- 1. A protein shown by SEQ ID NO: 1 or having the amino acid sequence derived therefrom in which one or more amino acids are substituted, deleted, or added.
 - 2. A protein encoded by a DNA hybridizing with the DNA shown by SEQ ID NO: 2.
 - 3. A DNA encoding the protein of Claim 1.

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- 4. A DNA hybridizing with the DNA shown by SEQ ID NO: 2.
- 5. A vector containing the DNA of Claim 3 or 4.
- 45 6. A transformant carrying the vector of Claim 5.
 - 7. A method of producing the protein of Claim 1 or 2, which comprises culturing the transformant of Claim 6.
 - 8. An antibody binding to the protein of Claim 1 or 2.

- 9. A method of screening a compound binding to the protein of Claim 1 or 2, which comprises a step of detecting the activity of the protein of Claim 1 or 2 to bind to a test sample.
- 10. A compound binding to the protein of Claim 1 or 2, wherein the compound have been isolated by the method of Claim 9.

Fig. 1

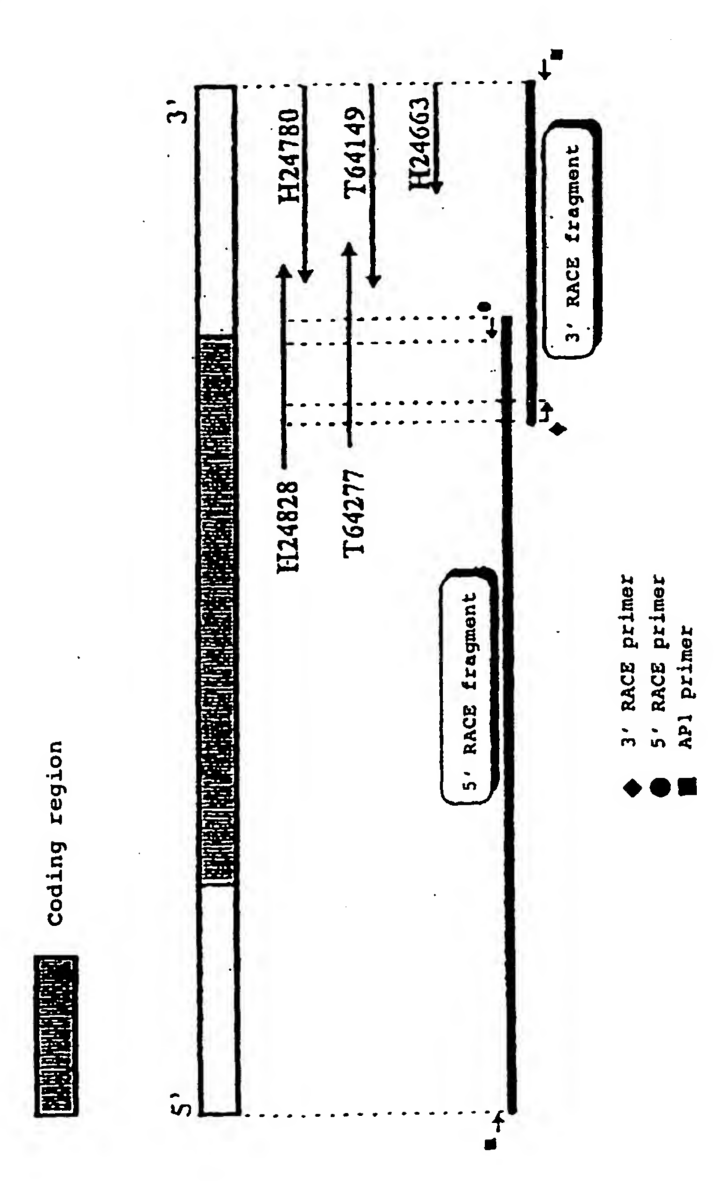


Fig. 2

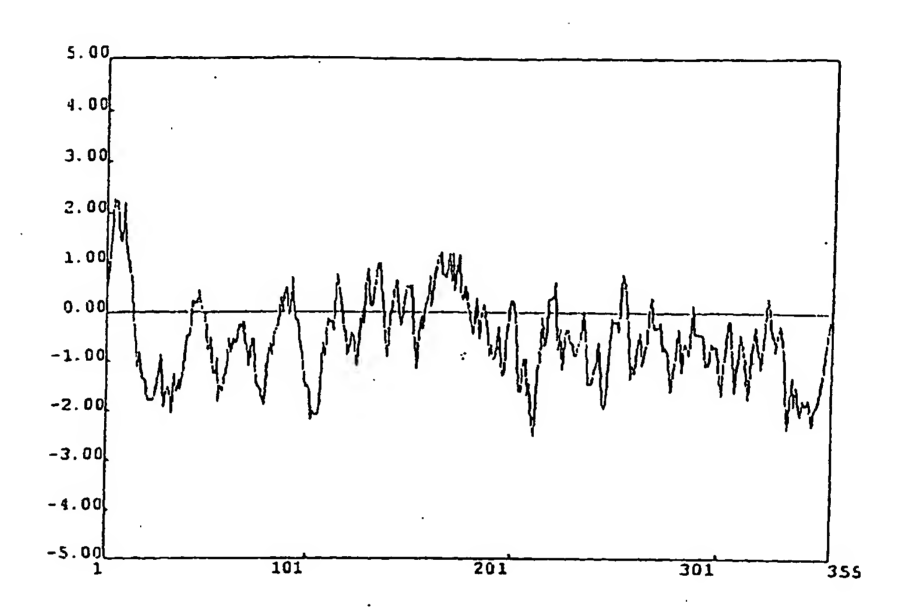
HSVEGFCC* H24828	MHLIGFFSYA CSLLAAALLP GPREAPAAAA AFESGLDISD AEPDAGEATA	50 50
HSVEGFCC H24828	YASKOLEEOL RSVSSVDELM TVLYPEYWKM YKCOLRKGGW QHNREQANLN	100
HSVEGFCC H24828	SRTEETIKFA AAHYNTEILK SIDNEWRKTQ CMPREVCIDV GKEFGVATNT	150 150
HSVEGFCC H24828	FFKPPCVSVY RCGGCCNSEG LOCMNTSTSY LSKTLFEITV PLSQGPKPVT	200 200
HSVEGFCC H24828	ISFANHTSCR CHSKLDVYRQ VHSIIRRSLP ATLPQCQAAN KTCPTNYMWN	250 250
HSVEGFCC H24828	NHICRCLAGE DFMFSSDAGD DSTDGFHDIC GPNKELDEET CQCVCRAGLR	300 300
HSVEGFCC H24828	PASCOPHKEL GRNSEGOVCK NKLFPSQCGA NREFDENTEQ CVCKRTEPRN	350 350
HSVEGFCC H24828	QPLNEGREAR ECTESPONCL LAGARETHOT ESCYREPETH ROKET-EPGF KLFHEDTESE E	400 400
HSVEGFCC H24828	SYSHEVERCY BSYWERHOMS	450 450
*HSVEGFCC:	human YEGF-C	

Fig. 3

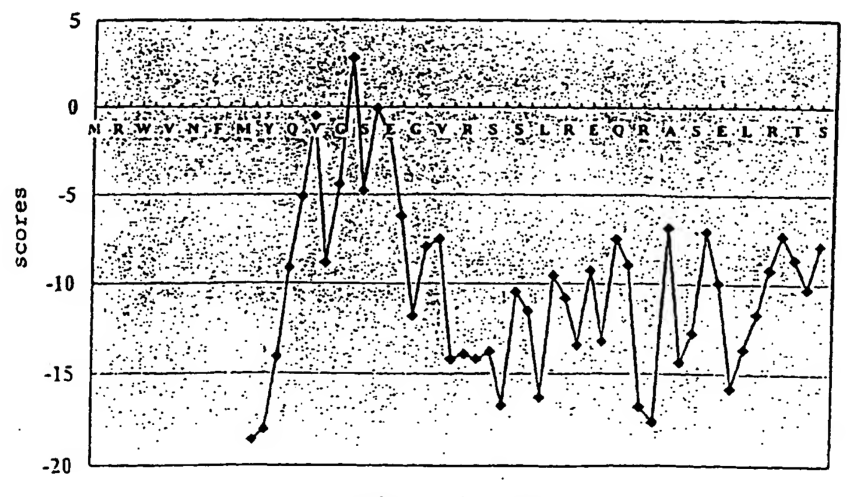
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	MHLUGFESVA C MRTUACULLU G MNRCWAUFLS L MPVMRLFPCF L	MMLYVOLVO SLUAAALIP CGYLAHVUA CCYLRLVSA OLLAGLAUP LALLLYUHH -LLLAALIO	GSSNEHGPVK GPREAPAAAA EEAEIPREVI EGDPIPEELY AVPPQQWALS AKWSQAAPMA LAPAQAPVSQ	EMLSD AGNGS EGGGQ	AEPDAGEATASQHS	50 50 50 50 50 50
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	YARKDLEEQU R IHRIRDLERU L	AASSLESLL SVSSVDSLM EIDSVGSED HGDPGEDG	RITHSEDANL TVLYPEYEMM S-L	WRERLELKSF YKGOLEKGGW	TSMDSRSASH QHNREQANLN DTSLRA DLNHTR	100 100 100 100 100 100
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	SRTEEÜIKFA P HGVHANKHVP E SHSGGELESL P	RGRRSLGSL SEVAVVP	VIDEEWORTO SIDNEWRKTO RSIEEAVPAV TOAEPAMIAE FOEV-WGRSY FMDV-YORSY WIEV-YTRAT	CSPRETCHEV CMPREVCIDV CKIPIVIYEI CKIBIEVEEI CRALERLUDV CHEIGTLUDI COPREVVUPL	ASELGKSINT GKEFGVAUNT PRSQVDPUSA SRRLIDRUNA VSEYPSEVEH FOEYPDEIEY TVELMGTVAK	150 150 150 150 150 150
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF-B	FFKPPCVS NELIWPPCVE NELVWPPCVE NELVWPPCVE NELVWPPCVE NELVWPPCVS LIFKFSCVP LIF	FROGGOCIE YROGGOCIS KROTGOCIT OROSGOCIN LROTGOCID MROGGOCID OROGGOCID	ESUI CMNUST EGLOCMNUST SSVKCOPSRV RNYOCRPTOV ENTHCVEVET EGLECVPTEE DGLEEVPTEE	SYISKELFEI SYLSKTLFEI HHRSVKVAKV OLRPVEVRKE ANVTMELKI SNITMEIMRI HQVRMEILMI	-SIPLTSVPE -TYPLSQGPK EYVEKKPKLK EIVEKKPIFK ESGDRPS KPHQGQH EYPSSQ-	200 200 200 200 200 200 200
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF-B	ENOVRLEERL E KATVTLEDEL A YMELTFSOEV R IGEMSFLOEN K	REMSKLDV RABATTSLN REEI-VAA	PRHPYSIIRE YRQVHSIIRE PDYREEDTGE ARPVTRSPGG LREKMKPERE RARQEKKSVE SA	SIQIPEEDEC S-LPATLPOC P-RESGKARK S-OEORAS R-PKGRGKER G-KGKGOFFK	OAANNTCPIN GBRLKPI. BEBORRI-	250 250 250 250 250 250 250
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	HLZDSHKEKE V YMZNNHIERE L			UDDD	CEETCOEVER TEODÁVTIÁT DEOTCKESEK DERTERERER	300 300 300 300 300 300 300
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	PALCEP AGLRPASCEP EN VRVRRPPKSK EN N-TDSRCKAR OI RRSFLRCOGR GE	RKFKHTHDK	TALBETOGA.			350 350 350 350 350 350
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	GCOKHKUFHE DI GPRNOPU-NE GP	GAGECTES	PQKCLLKGKK	OREPFHT FHHQTESCYR	RP CASGKTAC RPCTNRQKAC	400 400 400 400 400 400
HSVEGF-D HSVEGF-C HSPDGF-A HSPDGF-B HSPIGF2 HSVEGF HSVEGF-B	-EPGFSYSEE VC	RCVESYWE	REOMS		• • • • • • • • • • •	450 450 450 450 450 450

Fig. 4

a) Hydrophobicity



b) Prediction of the human VEGF-D signal peptide



amino acid sequence

INTERNATIONAL SEARCH REPORT International application No. PCT/JP97/02456 A. CLASSIFICATION OF SUBJECT MATTER Int. C16 C12N15/18, C12N15/63, C12P21/02, C07K14/485, C07K16/22, G01N33/50 According to International Patent Classification (IPC) or to both national classification and IPC FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) Int. C16 C12N15/18, C12N15/63, C12P21/02, C07K14/485, C07K16/22, G01N33/50 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WPI, WPI/L, BIOSIS PREVIEWS, CAS ONLINE, GENETYX-MAC/CD C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category* Relevant to claim No. Yamada, Y. et al. "Molecular cloning of a novel 1 - 10vascular endothelial growth factor, VEGF-D.* Genomics (1997, Jun.), Vol. 42, No. 3, p. 483-488 Vladimir, J. et al. "A novel vascular 1 - 2 endothelial growth factor, VEGF-C, (VEGFR-2) receptor tyrosine kinases" EMBO J. (1996, Jan.) Vol. 15, No. 2, p. 290-298 X Vladimir, J. et al. "A novel vascular 1 - 2 endothelial growth factor, VEGF-C, is a ligand for the Flt4(VEGFR-3) and KDR(VEGFR-2) receptor tyrosine kinases" EMBO J. (1996, Jan.) Vol. 15, No. 7, p. 1751 Maurizio, O. et al. "Identification of a c-fos-1 - 2 induced gene that is related to the plateletderived growth factor/vascular endothelial growth factor family" Proc. Natl. Acad. Sci. USA (1996, Oct.) Vol. 93, p. 11675-11680 Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents: later document published after the international filing date or priority document defining the general state of the art which is not considered date and not in conflict with the application but cited to understand the principle or theory underlying the invention to be of particular relevance "E" earlier document but published on or after the international filing date document of particular relevance; the claimed invention cannot be document which may throw doubts on priority claim(s) or which is considered novel or cannot be considered to involve an inventive cited to establish the publication date of another citation or other step when the document is taken alone special reason (as specified) document of particular relevance; the claimed invention cannot be "O" document referring to an oral disclosure, use, exhibition or other considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art document published prior to the international filling date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report October 7, 1997 (07. 10. 97) October 21, 1997 (21. 10. 97) Name and mailing address of the ISA/ Authorized officer Japanese Patent Office Facsimile No. Telephone No. Form PCT/ISA/210 (second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP97/02456

		FC1/3	P97/02456
C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT		
lategory*	Citation of document, with indication, where appropriate, of the relev	vant passages	Relevant to claim No
X	Georg. B. et al. "Expression of vascular endothelial growth factor during embryongiogenesis and endothelial cell differentiation" Development (1992) Vo. p. 521-532	onic	1 - 10
X	David, T.S. et al. "The mouse gene for endothelial growth factor" J. Biol. Che (1996, Feb.) Vol. 271, No. 7, p. 3877-	em.	1 - 10
x	Kevin, P.C. et al. "Vascular endothelia factor" J. Biol. Chem. (1992) Vol. 267 p. 16317-16322	al growth, No. 23,	1 - 10
х	Greg, C. et al. "Amino acid and cDNA so of a vascular endothelial cell mitogen homologous to platelet-derived growth Proc. Natl. Acad. Sci. USA (1990) Vol. p. 2628-2632	that is factor"	1 - 10
х	Edmund, T. et al. "The human gene for endothelial growth factor" J. Biol. Che Vol. 266, No. 18, p. 11947-11954	vascular em. (1991)	1 - 10
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INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP97/02456

Disclosure other than written disclosures

- 1. The GenBank Database (Rel. 100) on GENETYX, Accession No. D89628, Yoshiki Yamada, Chugai Research Institute for Molecular Medicine. (29-Nov-1996)
- 2. The GenBank Database (Rel. 100) on GENETYX, Accession No. T64277, Hillier, L. et al. (1995)

Form PCT/ISA/210 (extra sheet) (July 1992)